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### Authors

Navarro-Mendoza, María Isabel  
Pérez-Arques, Carlos  
Panchal, Shweta  
et al.

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**Early diverging fungus *Mucor circinelloides* lacks  
centromeric histone CENP-A and displays a mosaic of point and  
regional centromeres**

María Isabel Navarro-Mendoza<sup>1#</sup>, Carlos Pérez-Arques<sup>1#</sup>, Shweta  
Panchal<sup>2</sup>, Francisco E. Nicolás<sup>1</sup>, Stephen J. Mondo<sup>3,4</sup>, Promit Ganguly<sup>2</sup>,  
Jasmyn Pangilinan<sup>3</sup>, Igor V. Grigoriev<sup>3,5</sup>, Joseph Heitman<sup>6\*</sup>, Kaustuv  
Sanyal<sup>2\*</sup>, and Victoriano Garre<sup>1\*</sup>

<sup>1</sup>Department of Genetics and Microbiology, Faculty of Biology, University  
of Murcia, Murcia, 30100, Spain; <sup>2</sup>Molecular Biology and Genetics Unit,  
Jawaharlal Nehru Centre for Advanced Scientific Research, Jakkur,  
Bangalore 560064, India; <sup>3</sup>US Department of Energy Joint Genome  
Institute, Walnut Creek, California, USA; <sup>4</sup>Bioagricultural Science and Pest  
Management Department, Colorado State University, Fort Collins, CO  
80521, USA; <sup>5</sup>Department of Plant and Microbial Biology, University of  
California Berkeley, Berkeley, CA 94598, USA; <sup>6</sup>Department of Molecular  
Genetics and Microbiology, Duke University Medical Center, NC, USA

#Equally contributed; \*Corresponding authors

Author Contact: Victoriano Garre, [vgarre@um.es](mailto:vgarre@um.es); Kaustuv Sanyal,  
[sanyal@jncasr.ac.in](mailto:sanyal@jncasr.ac.in); Joseph Heitman, [heitm001@duke.edu](mailto:heitm001@duke.edu).

Lead Contact: Victoriano Garre, [vgarre@um.es](mailto:vgarre@um.es)

**Summary**

Centromeres are rapidly evolving across eukaryotes, despite  
performing a conserved function to ensure high fidelity chromosome

24 segregation. CENP-A chromatin is a hallmark of a functional centromere in  
25 most organisms. Due to its critical role in kinetochore architecture, the  
26 loss of CENP-A is tolerated in only a few organisms, many of which  
27 possess holocentric chromosomes. Here, we characterize the  
28 consequence of the loss of CENP-A in the fungal kingdom. *Mucor*  
29 *circinelloides*, an opportunistic human pathogen, lacks CENP-A along with  
30 the evolutionarily conserved CENP-C, but assembles a monocentric  
31 chromosome with a localized kinetochore complex throughout the cell  
32 cycle. Mis12 and Dsn1, two conserved kinetochore proteins were found to  
33 colocalize to a short region, one in each of nine large scaffolds, comprised  
34 of an ~200-bp AT-rich sequence followed by a centromere-specific  
35 conserved motif that echoes the structure of budding yeast point  
36 centromeres. Resembling fungal regional centromeres, these core  
37 centromere regions are embedded in large genomic expanses devoid of  
38 genes yet marked by Grem-LINE1s, a novel retrotransposable element  
39 silenced by the Dicer-dependent RNAi pathway. Our results suggest that  
40 these hybrid features of point and regional centromeres arose from the  
41 absence of CENP-A, thus defining novel mosaic centromeres in this early-  
42 diverging fungus.

## 43 **Introduction**

44 Accurate chromosome segregation is crucial to maintain genome  
45 integrity during cell division. The timely attachment of microtubules to  
46 centromere DNA is essential to achieve proper chromosome segregation.  
47 This is accomplished by a specialized multilayered protein complex, the

48 kinetochore which links microtubules to centromere DNA. This protein  
49 bridge is divided into two layers – the inner and outer kinetochore. The  
50 fundamental inner kinetochore protein is the histone H3 variant CENP-A. It  
51 binds directly to centromere DNA and lays the foundation to recruit other  
52 essential proteins of the kinetochore complex, playing a fundamental role  
53 in centromere structure and function, and hence, precise chromosome  
54 segregation [1,2]. CENP-A is also found at all identified neocentromeres  
55 [3] and at the active centromeres of dicentric chromosomes [4], acting as  
56 an epigenetic determinant of centromeric identity.

57         Despite its conserved function, the centromere is one of the most  
58 rapidly evolving regions of the genome [5]. This so-called “centromere  
59 paradox” has led to centromeres of diverse sizes and content. The first  
60 centromeres identified in *Saccharomyces cerevisiae* were found to be  
61 point centromeres - small regions of ~120 bp defined by specific DNA  
62 sequences [6,7]. In contrast to point centromeres described in only a few  
63 budding yeasts of the phylum Ascomycota, most other fungi and  
64 metazoans have regional centromeres that are larger, ranging from a few  
65 kilobases to several megabases [8]. Regional centromeres are often  
66 interspersed with repetitive sequences and are mostly defined by  
67 epigenetic factors rather than DNA sequence per se [9]. While the  
68 centromeres described thus far are confined to one region of the DNA,  
69 there are organisms in which their entire chromosomes are loaded with  
70 kinetochores and display a parallel separation of the two chromatids. Such  
71 chromosomes possess holocentromeres and have been found in several  
72 clades of insects [10] and nematodes like *Caenorhabditis elegans* [11].

73 Despite this remarkable variation in the types of centromeres, most  
74 organisms have their centromere loci defined by the binding of CENP-A.  
75 However, there are a few exceptions, including some insect lineages and  
76 kinetoplastids, which have independently lost CENP-A. The kinetoplastid  
77 *Trypanosoma brucei* has evolved a unique set of proteins to perform the  
78 role of the kinetochore complex [12], whereas all insects that have  
79 recurrently lost CENP-A have transitioned from monocentricity to  
80 holocentricity [10].

81         Neither holocentricity nor lack of CENP-A has ever been reported in  
82 the fungal kingdom until a recent study showed that this protein could be  
83 absent in the Mucoromycotina subphylum [13], which belongs to the early  
84 diverging fungi. Kinetochore structure and centromere function are well-  
85 studied in species of the Dikarya, whereas centromere identity as well as  
86 the kinetochore architecture remain largely unexplored in basal fungi  
87 because these are challenging to manipulate genetically. *Mucor*  
88 *circinelloides*, a fungus of the subphylum Mucoromycotina, is an exception  
89 and molecular tools are available to modify its genome [14]. *M.*  
90 *circinelloides*, as with other Mucorales, causes an infectious disease  
91 known as mucormycosis, which have been associated with high mortality  
92 rates due to rhino-orbital-cerebral, pulmonary, or cutaneous infections  
93 [15]. Despite its medical significance and research efforts to identify new  
94 antifungal targets, the genome biology and cell cycle of this fungus  
95 remain poorly understood. Using evolutionarily conserved kinetochore  
96 proteins as tools, we investigated the dynamics of the kinetochore  
97 complex during nuclear division and unveiled the unique characteristics of

98 the centromeres of *M. circinelloides*. Studying distinct and divergent  
99 factors involved in the cell cycle of this organism will open avenues for the  
100 development of species-specific antifungal drugs.

## 101 **Results**

### 102 **Last Mucoralean common ancestor lost CENP-A and CENP-C, and** 103 **the descendant Mucoromycotina lack centromere-specific histone** 104 **variants**

105 A recent study of kinetochore evolution in eukaryotes reported the  
106 striking absence of two well conserved inner kinetochore proteins, CENP-A  
107 and CENP-C, in two species of the Mucoromycotina [13]. To ascertain the  
108 presence or loss of well-studied kinetochore proteins across the entire  
109 subphylum, we analyzed the curated proteomes of 75 fungal species,  
110 including 55 species of Mucoromycotina and 20 species from other fungal  
111 clades (Table S1). The kinetochore protein sequences from *Mus musculus*,  
112 *Ustilago maydis*, *S. cerevisiae*, and *Schizosaccharomyces pombe* served  
113 as queries to conduct iterative Blast and Hmmer searches against these  
114 proteomes. These sequences were confirmed as putative orthologs by the  
115 presence of conserved Pfam domains or by obtaining a matching first hit  
116 in a reciprocal Blastp search against the initial four species: mouse, *U.*  
117 *maydis*, budding, and fission yeasts (Table S1, protein sequences at  
118 Mendeley data DOI: 10.17632/wpyyb58h8v.1).

119 To confirm the presence or absence of centromere-specific histone  
120 H3 CENP-A in each fungal species, 289 histone H3 homologs were  
121 analyzed, identifying 27 putative CENP-A proteins (Figure S1, Table S1,

122 protein sequences at Mendeley data DOI: 10.17632/wpyyb58h8v.1). These  
123 proteins share common features with well-known CENP-A proteins [16],  
124 such as an N-terminal tail that varies significantly in length and sequence  
125 compared to the canonical H3 counterparts (Figure S1B), the absence of a  
126 conserved glutamine residue in the first  $\alpha$ -helix (69Q) and a phenylalanine  
127 residue (84F) (Figure S1C), a longer and divergent first loop (Figure S1C),  
128 and a non-conserved C-terminal tail (Figure S1D). In addition, the histone  
129 fold domain (HFD) sequences of the putative CENP-A proteins have  
130 diverged rapidly from the conserved canonical H3 sequences [16] (Figure  
131 S1E). Nine additional H3 homologs also share several CENP-A common  
132 features but were regarded as rare histones because they lack these  
133 conserved substitutions. The analysis revealed that CENP-A was absent in  
134 most of the Mucoromycotina clade, specifically in the orders  
135 Umbelopsidales and Mucorales; nevertheless, we identified putative CENP-  
136 A proteins in all Endogonalean species (Figure 1). Similarly, CENP-C is  
137 present in the Endogonales and Umbelopsidales but not in the Mucorales  
138 (Figure 1). These findings suggest that the last Mucoromycotina common  
139 ancestor possessed CENP-A, which was lost after the Endogonales  
140 diverged from the rest of the clade. In the absence of CENP-A, CENP-C was  
141 presumably lost soon after. In addition, CENP-Q is absent in all Mucoralean  
142 species, and CENP-U is lost in the whole subphylum. These proteins have  
143 been recurrently lost across the fungal kingdom, as evidenced by their  
144 absence in species of all phyla of early-diverging fungi and most species  
145 in Basidiomycota (Figure 1). The absence of orthologs that were present in  
146 closely-related species should be taken with caution because this could be

147 a result of the incomplete state of their genome assemblies and  
148 annotation.

149         Despite these relevant losses, the remaining kinetochore proteins  
150 were well conserved among most species of Mucoromycotina, especially  
151 in *M. circinelloides*. Therefore, we hypothesized that another histone  
152 protein could have taken the role of CENP-A as a part of centromeric  
153 nucleosome for kinetochore assembly. We searched the *M. circinelloides*  
154 genome sequence for histone protein-coding genes and examined their  
155 sequences seeking distinctive features of histone variants. Genes  
156 encoding mammalian histone variants have intronic sequences and  
157 polyadenylation [poly(A)] signals, in contrast to canonical histones H3.1  
158 and H3.2 [17]. However, fungal canonical histones may have intronic  
159 sequences and poly(A) signals [18]. Thus, the analysis of possible variants  
160 focused on the presence of substitutions in relevant amino acid residues  
161 targeted by post-translational modifications [19] rather than intronic and  
162 poly(A) sequences. The *M. circinelloides* genome contains four histone H3-  
163 and three histone H4-coding genes, named *histone H three* (*hht1-4*) and  
164 *histone H four* (*hhf1-3*), respectively (Figure S2A), and all of them feature  
165 a putative poly(A) signal (5'-AATAAA-3'). Three of the four histone H3-  
166 coding genes are intronless and their protein sequence identical, except  
167 the intron-containing gene *hht4* (Figure S2A). Also, the amino acid  
168 residues at positions 32, 43, and 97 in the Hht4 protein sequence differ  
169 compared to the conserved H3 sequence (Figure S2B). Similarly, histone  
170 H4-coding genes *hhf2* and *hhf3* lack introns and encode identical proteins,



171 whereas *hhf1* has intronic sequences (Figure S2A) and its product has a  
172 longer N-terminal tail (Figure S2C).

173         To test if the minor differences observed in Hht4 and Hhf1 could  
174 have resulted in a centromere-specific binding protein, the cellular  
175 localization was analyzed and compared to their canonical histone  
176 counterpart Hhf3. We constructed alleles that contained each histone-  
177 coding gene fused in-frame at the C-terminus with the enhanced green  
178 fluorescent protein (eGFP) flanked by each of the histone gene native  
179 promoter and terminator sequences, and a selectable marker (Figure S3).  
180 Homokaryotic strains expressing fluorescent histones Hht4, Hhf1, or Hhf3  
181 were obtained by transforming a wild-type strain with their respective  
182 eGFP-tagged alleles. Integration by homologous recombination at the  
183 corresponding native loci was confirmed by PCR (Figure S3 and Table S2).

184         *M. circinelloides* is a saprophytic fungus with defined sexual and  
185 asexual lifecycles [20]. The asexual reproductive cycle is the preferred  
186 culture method to maintain this organism under laboratory conditions, due  
187 to fast developing mycelia and sporangiospores that contain asexual  
188 spores [14]. Asexual spores from these strains were observed with a  
189 confocal microscope under two stages of germination: ungerminated and  
190 4-h pre-germinated spores, that had started swelling and forming germ  
191 tubes. In both conditions, all fluorescent histone fusions were localized  
192 encompassing the entire nucleus instead of forming kinetochore-like  
193 nuclear clusters (Figure S2D). Most species of the Mucoromycotina are  
194 coenocytic fungi and have multinucleated spores, as does the *M.*

195 *circinelloides* wild-type strain used in this study which has large spores  
196 with a high number of nuclei [21]. We established that nuclear division in  
197 *M. circinelloides* is asynchronous by observing the nuclear distribution of  
198 Hht4-eGFP protein in pre-germinated spores during a time-lapse imaging  
199 session (Video S1). Altogether, these findings indicate that either none of  
200 these histone variants functions like CENP-A in kinetochore assembly or  
201 *Mucor* centromeres are holocentric in nature. We could test these two  
202 possibilities by studying other kinetochore proteins that are evolutionarily  
203 conserved and easily identifiable in *M. circinelloides*.

#### 204 **Mis12 and CENP-T complexes show discrete nuclear localization**

205 Analyzing the nuclear localization and assembly dynamics of the  
206 kinetochore complex of *M. circinelloides* could offer insights into its  
207 architecture. In the absence of CENP-A or another similar histone H3  
208 variant, and CENP-C, that could anchor the kinetochore formation, we  
209 focused our attention on the evolutionarily conserved and obvious  
210 homologs of inner kinetochore proteins that are expected to be closer to  
211 the centromere DNA in this organism (Figure 2A). We designed  
212 fluorescent kinetochore fusion proteins of Mis12, Dsn1, and CENP-T  
213 because these are well-conserved in almost all species of the  
214 Mucoromycotina. First, we constructed alleles to tag the Mis12 and Dsn1  
215 proteins with the red fluorescent protein mCherry, both at their C-termini  
216 and N-termini, aiming to integrate the alleles in their endogenous loci  
217 following a similar strategy as for the epitope tagging of the histone genes  
218 described above. Homologous recombination in the desired transformants

219 was confirmed by PCR, and six colonies from the four tagged alleles were  
220 checked for fluorescent localization. Unfortunately, fluorescent signals  
221 were not detected in any of the four different tagged strains possibly due  
222 to low levels of expression.

223 As an alternative approach, the Mis12 and Dsn1 proteins were  
224 tagged with mCherry at each end, N-terminal and C-terminal, separately,  
225 and overexpressed from a strong promoter (*Pzrt1*) in *M. circinelloides*  
226 (Figures 2B and S3). The alleles were designed to target integration into  
227 the *carRP* locus, a gene encoding an enzyme involved in carotenogenesis.  
228 Thus, the mutant strains obtained should lack colored-carotenes and  
229 display an albino phenotype [14]. After confirming the integration of the  
230 alleles in the *carRP* locus by PCR (Figures S3E and S3F, Table S2), one  
231 strain harboring each construct was used as a parental strain to integrate  
232 the allele for H3-eGFP expression at its own locus as previously described,  
233 allowing monitoring colocalization of the kinetochore proteins within the  
234 nucleus. Once the double integration was confirmed, two independent  
235 strains of each type were selected for fluorescent localization.

236 Mis12 and Dsn1 displayed a clear localization signal forming small  
237 fluorescent puncta within the nucleus marked by histone H3 (Figures 2C  
238 and 2D, respectively). The microscopy screening confirmed that the  
239 expression of both N-terminal and C-terminal tagged proteins was similar.  
240 The Mis12 and Dsn1 localization patterns revealed a single cluster of  
241 kinetochores as observed in many fungal species. This result strongly

242 suggests that *M. circinelloides* has monocentric chromosomes with a  
243 localized kinetochore even in the absence of CENP-A.

244       Next, another evolutionarily conserved inner kinetochore protein  
245 CENP-T, which is known to interact with DNA, was tagged with eGFP at the  
246 C-terminus following the same procedure used for the histone tagging and  
247 integrated by homologous recombination at the endogenous locus in  
248 strains expressing the Mis12-mCherry and Dsn1-mCherry fusion proteins.  
249 Integration of the alleles at the desired loci was confirmed by PCR in both  
250 parental backgrounds (Figure S3D), and the localization of the kinetochore  
251 proteins was examined by analyzing their fluorescent signal in pre-  
252 germinated spores. CENP-T colocalized with Mis12 and Dsn1 in each  
253 nucleus (Figures 2E and 2F, respectively), indicating that all three proteins  
254 assemble together in the kinetochore ensemble. Time-lapse imaging  
255 during spore germination of the double-tagged strains allowed the  
256 analysis of nuclear division and kinetochore structure during the cell cycle  
257 in live cells. The Mis12 and Dsn1 proteins clustered at the nuclear  
258 periphery during division (Figure 2G), a typical feature of fungal  
259 kinetochore proteins, suggesting all these three proteins are constitutively  
260 present at the kinetochore during the cell cycle.

## 261 ***M. circinelloides* exhibits small mosaic centromeres associated** 262 **with RNAi-suppressed L1-like retrotransposons**

263       The centromere regions of early-diverging fungi have never been  
264 described, possibly because of their genetic intractability. The generation  
265 of kinetochore-tagged strains of *M. circinelloides* offered the opportunity

266 to identify the nature of centromeres in an organism that lost CENP-A and  
267 CENP-C. To map *M. circinelloides* centromeres, chromatin  
268 immunoprecipitation followed by next generation sequencing (ChIP-seq)  
269 was performed with Mis12- and Dsn1-mCherry proteins. Illumina  
270 sequencing was conducted in two independently immunoprecipitated (*IP*  
271 DNA) samples for each kinetochore protein, one for Mis12-mCherry and  
272 the other for Dsn1-mCherry, using RFP-Trap MA beads (Chromotek) for  
273 mCherry immunoprecipitation. Input control samples consisting of total  
274 sheared DNA (*Input* DNA) were also sequenced for each tagged strain, as  
275 well as mock binding control (*beads only* DNA) samples with MA beads  
276 without antibody. Raw sequencing data were processed, and reads were  
277 aligned to a newly generated PacBio genome assembly of *Mucor*  
278 *circinelloides* f. *lusitanicus* MU402 strain (available at JGI  
279 [http://genome.jgi.doe.gov/Muccir1\\_3/Muccir1\\_3.home.html](http://genome.jgi.doe.gov/Muccir1_3/Muccir1_3.home.html), hereafter the  
280 *M. circinelloides* genome), the parental strain of the kinetochore protein-  
281 tagged strains. Out of the whole genome assembly of 24 scaffolds, which  
282 has an average GC content of 42.19%, 11 major scaffolds encompass  
283 more than 90% of the genome.

284 Both Mis12 and Dsn1 *IP* DNA sequence reads were compared to the  
285 corresponding control *Input* DNA reads to identify statistically significant  
286 kinetochore protein-enriched genomic loci in both *IP* DNA samples ( $\text{FDR} \leq$   
287  $5 \times 10^{-5}$ , fold enrichment  $\geq 1.6$ ; Table S3). The result of this analysis was  
288 visualized in a genome browser to ensure that the enrichment was robust  
289 and absent in either the *Input* or *beads only* DNA controls (Figure S4). By  
290 this approach, we detected nine significantly enriched peaks that

291 overlapped in both Mis12 and Dsn1 *IP* DNA samples, and thus we defined  
292 kinetochore protein-bound regions as core centromeres (CCs) (Figure 3A)  
293 that were designated by the corresponding scaffold number. Each of these  
294 core *CEN* sequences was located in a different major scaffold of the  
295 genome assembly (Figure 3B), indicating that each chromosome was  
296 assembled into an average of one to three scaffolds. The identification of  
297 five repeated regions matching the telomeric sequence (5'-TTAGGG-3') at  
298 the 3'-end of scaffolds 2, 3, 8, 9, and 10, suggested that CC2 and CC9  
299 display a subtelomeric localization (Figure 3B). Because the length of the  
300 overlapping peaks was minimally different in the two kinetochore *IP* DNA  
301 samples, DNA sequence enriched with at least one of the kinetochore  
302 proteins was assumed to be the length of kinetochore binding region on  
303 each chromosome. In addition, contiguous enriched regions were added to  
304 each CC, defining the core centromere regions (Table S3).

305       The average length of core centromere regions of *M. circinelloides* is  
306 941 bp (Figure 4A). These unusually short regions are consistently AT-rich  
307 (71% average) (Figure 4B), resembling point centromeres of *S. cerevisiae*  
308 and its related budding yeast species. Further analysis revealed a 41-bp  
309 DNA sequence motif conserved in all nine kinetochore protein-bound core  
310 centromere regions (Figure 4C). An extensive search of the presence of  
311 this motif across the whole genome of *M. circinelloides* revealed that the  
312 41-bp sequence is centromere-specific, occurs as a single-copy element at  
313 each core centromere, and is absent in the rest of the genome. Moreover,  
314 a closer inspection of the core centromere regions revealed a conserved  
315 pattern. Each core centromere is composed of a minor peak enriched of

316 kinetochore proteins spaced by a highly AT-rich stretch spanning the  
317 major binding peak of kinetochore proteins that starts with the 41-bp  
318 conserved motif (Figure 4D).

319       Surrounding the core centromere regions, both upstream and  
320 downstream, are stretches of genomic sequences completely devoid of  
321 genes. These pericentric regions are considerably larger than the core and  
322 vary in length, ranging from 15 to 75 kb (Figure 5A). These pericentric  
323 regions contain a variable number of large repeats oriented away from the  
324 core centromere regions (Figure S5). These repeats cluster in nine groups  
325 according to their similarity ( $\geq 95\%$  identity and  $\geq 70\%$  coverage across  
326 their entire sequence). Clusters 1, 2 and 3 contain all of the full-length  
327 elements, which were termed autonomous elements, while the remaining  
328 clusters are composed of incomplete repeats or remnants (Figure S6A). A  
329 pairwise comparison of all of the repeats revealed a high similarity in their  
330 aligned sequences ( $\geq 90\%$  identity), though the truncated sequences of  
331 remnants showed higher differences accounting for the gaps in the  
332 alignment.

333       Each autonomous repeat comprises two sequential open reading  
334 frames (ORFs) with an overlapping codon. The first ORF encodes a product  
335 containing several zinc finger domains, and the second codes for a protein  
336 with AP endonuclease and reverse transcriptase domains (Figure 5B,  
337 Table S4). Based on their high similarity, their ORF domain architecture,  
338 the presence of a poly(A) signal at their 3'-end, and the absence of long  
339 terminal repeats (LTR) at either end, they were classified as a single non-

340 LTR retrotransposon. Phylogenetic profiling revealed that these  
341 retrotransposons belong to the LINE1 clade (Figure S6B) and are closely  
342 related to other LINE1-like retrotransposons of Mucoromycotina species,  
343 so they were termed Genomic Retrotransposable Element of  
344 Mucoromycotina (Grem) LINE1-like. A search of the *M. circinelloides*  
345 genome revealed that the Grem-LINE1s are found specifically in the  
346 pericentromeric regions; or at the non-telomeric ends of three small  
347 scaffolds, oriented away from the end (Table S4). Four major scaffolds end  
348 abruptly in a putative centromere region, suggesting that they may be  
349 linked to these minor scaffolds that contain the Grem-LINE1 elements at  
350 one end. Broadening the search across 55 genomes of species of  
351 Mucoromycotina identified the Grem-LINE1 elements in most Mucorales  
352 and Umbelopsidales ( $\geq 30\%$  identity and  $\geq 50\%$  coverage), but they could  
353 not be found in any of the Endogonalean genomes, indicating an inverse  
354 correlation between the presence of Grem-LINE1s and CENP-A (Figure 1  
355 and Table S5).

356 RNAi silences expression and prevents transposition of  
357 retrotransposable elements in several fungal species featuring functional  
358 RNAi [22,23]. *M. circinelloides* possess an elaborate RNAi mechanism that  
359 protects the genome against invading elements, that could prevent the  
360 movement of retrotransposable elements. The two Dicer enzymes,  
361 particularly Dcl2, and one of the three Argonaute proteins (Ago1), play a  
362 pivotal role in this protective RNAi-related pathway [24]. To analyze the  
363 transcriptional landscape of the pericentric regions, previously generated  
364 and publicly available transcriptomic raw data for mRNA and small RNAs



365 were reprocessed and aligned to the *M. circinelloides* genome assembly.  
366 The pericentric regions were almost devoid of transcription in the wild-  
367 type strain, although the retrotransposons were modestly expressed.  
368 Indeed, a high number of sRNAs corresponding to the retrotransposons  
369 were detected (Figures 5C and S6), indicating an inverse correlation  
370 between mRNA and sRNA levels. This inverse correlation was confirmed  
371 by analyzing the transcriptomic profile of mutant strains lacking essential  
372 components of the RNAi machinery, Ago1 or Dcl1/Dcl2. In these RNAi-  
373 deficient mutants, the pattern was reversed; high mRNA levels from the  
374 retrotransposons were observed, while the production of sRNAs  
375 originating from those regions was considerably lower than in the wild-  
376 type strain. These findings suggest that the retrotransposons contained in  
377 the pericentric regions of *M. circinelloides* are silenced by the Dicer-  
378 dependent RNAi pathway.

379         In most organisms, deposition of CENP-A at the centromere-specific  
380 nucleosomes is correlated to depletion of histone H3 [25-27]. Because  
381 CENP-A is absent in *M. circinelloides*, we quantified the presence of  
382 histone H3 at the centromere-specific nucleosomes by performing ChIP  
383 with anti-histone H3 antibodies and a qPCR analysis. The occupancy of  
384 histone H3 was examined by qPCR in four regions across the scaffold\_2:  
385 the spanning kinetochore protein-binding region (core centromere, CC2),  
386 the ORF-free pericentromeric region, the centromere-flanking ORFs and  
387 the control region far-CEN ORF ~2 Mb away from the centromere (Figure  
388 5D). There are no significant differences in histone H3 occupancy across  
389 the analyzed regions, indicating that H3 is not depleted at either the

390 kinetochore protein binding region or the pericentric region compared to  
391 the flanking ORFs and the control far-CEN ORF (Figure 5D).

## 392 **Discussion**

393       Mucoralean species are the causative agents of mucormycosis, an  
394 emerging fungal infection with extremely poor clinical outcomes, probably  
395 as a consequence of their innate resistance to all current antifungal drugs  
396 [28,29] and their ability to evade host innate immunity [30,31]. In spite of  
397 their clinical relevance, limited information is available on essential  
398 biological processes in these fungal pathogens, especially nuclear division  
399 and cell cycle. In this study, we identified major kinetochore components  
400 in *M. circinelloides*, and utilized them to identify and characterize nine  
401 chromosomal loci as centromeres.

402       In most eukaryotes, the centrochromatin is composed of  
403 nucleosomes containing the histone H3 variant CENP-A as a hallmark of  
404 centromere identity. The most striking observation of our study was the  
405 absence of this centromere determining protein in all Mucoralean species.  
406 Absence of CENP-A was accompanied by loss of CENP-C, while most other  
407 kinetochore protein orthologs were present in these species. Loss of a  
408 previously-thought indispensable protein like CENP-A raised questions on  
409 the evolutionary bases of centromere and kinetochore structure. Absence  
410 of CENP-A had been previously observed in certain insect orders including  
411 butterflies and moths, bugs and lice, earwigs, and dragonflies which in all  
412 known cases led to a transition to holocentricity, suggesting that the

413 presence of a centromere-specific protein is dispensable for segregation  
414 of a chromosome with localized kinetochore[10].

415 Our study is the first to demonstrate loss of core kinetochore  
416 proteins in the fungal kingdom, reinforcing that the absence of CENP-A  
417 can be tolerated in eukaryotic organisms. Surprisingly, punctate  
418 localization patterns of other conserved kinetochore proteins including  
419 Mis12, Dsn1, and CENP-T throughout the cell cycle indicate that these  
420 organisms still retain a monocentric arrangement of a canonical  
421 kinetochore structure, as opposed to the holocentric insects that lack  
422 CENP-A. Kinetochore proteins in *M. circinelloides* were observed to be  
423 localized at the periphery of each nucleus in the asexual spores as well as  
424 mycelia. Three types of nuclear divisions have been studied in  
425 multinucleated fungi – synchronous, asynchronous, and parasynchronous  
426 [32]. In all ascomycetes except *S. pombe* and *Zymoseptoria tritici*,  
427 kinetochore proteins are centromere-localized and clustered throughout  
428 the cell cycle [33,34]. In the basidiomycete *C. neoformans*, kinetochore  
429 proteins transition between clustering and declustering states in different  
430 stages of the cell cycle [35].

431 We have observed asynchronous nuclear divisions during  
432 germination of asexual spores of *M. circinelloides*, a multinucleated and  
433 coenocytic fungus. The kinetochore was found to be consistently present  
434 as a cluster in all stages of mitosis, implying that the basic kinetochore  
435 structure in this organism is constitutively assembled throughout the cell  
436 cycle. Overall, these findings hint that the role of CENP-A is being carried

437 out by either a distinct protein that has a newly evolved function or by  
438 one of the known kinetochore proteins like CENP-T, which also has a  
439 histone-fold DNA binding domain. We also tried to find whether one of the  
440 histone H3 variants plays a role of centromeric histone, but none of them  
441 displayed kinetochore-like localization, indicating that they are probably  
442 not involved in replacing CENP-A function in chromosome segregation  
443 mediated by the kinetochore machinery. Usually, histone H3 is depleted  
444 at the centromeres as compared to other non-centromeric loci in the  
445 genome. We observed the presence of H3 at the relatively short  
446 kinetochore protein-binding region at a level that is similar to the  
447 pericentric region or gene bodies. Thus, we propose that centromeres in  
448 this organism contain histone H3 at the centromere core. Further studies  
449 will provide the exact composition of centromere-specific nucleosomes in  
450 this basal fungus.

451         A ChIP-based analysis using Dsn1 and Mis12 revealed the first  
452 glimpse of the centromere structure of this *Mucoromycotina* species. The  
453 centromeres in this organism possess features of both point and regional  
454 centromeres, resulting in a unique mosaic; thus, we named them mosaic  
455 centromeres (Figure 6). One of their distinctive features is a small  
456 kinetochore protein binding domain (~750-1150 bp) defined by the  
457 presence of an AT-rich sequence stretch (~70-80% AT, ~100 bp), which is  
458 followed by a conserved centromere-specific 41-bp sequence motif. This  
459 motif is only found in *M. circinelloides* centromeres and marks the start of  
460 the major kinetochore-binding region. Overall, this pattern resembles the  
461 distribution of conserved elements in *S. cerevisiae* point centromeres, in

462 which two conserved Centromere Determining Elements (CDEI and CDEIII)  
463 are separated by an AT-rich non-conserved DNA sequence stretch (CDEII)  
464 [7]. The kinetochore of *S. cerevisiae* binds CDEI and CDEIII in a sequence-  
465 dependent manner through Cbf1 [36] and the CBF3 complex [37,38],  
466 respectively. The presence of a conserved DNA motif in the centromeres  
467 of *M. circinelloides* suggests that it could be a binding site for an unknown  
468 centromere-specific protein in this species.

469 Other features of *M. circinelloides* mosaic centromeres are  
470 reminiscent of fungal regional centromeres like the ones described in  
471 several *Candida* spp. [39,40]. The pericentric regions of *M. circinelloides*  
472 are large (~15-75 kb), gene-free, and transcriptionally silenced sequences  
473 that are interspersed by Grem-LINE1s, which are repeats of a LINE1-like  
474 non-LTR retrotransposable element. Several retroelements have been  
475 identified in the centromeres and neocentromeres of highly diverged  
476 eukaryotes: non-LTR LINE1-like elements in mammals [41-46] and *Jockey*  
477 elements in the fruit fly [47], and LTR retrotransposons in plants [48] and  
478 fungi [23,49]. Based on these findings, a model has been proposed that  
479 CENP-A is recruited by genomic sequences rich in repeats and  
480 transposable elements and thus, gives rise to the centromeres [50]. The  
481 Grem-LINE1s are arranged in a systematic pattern, oriented away from  
482 the core centromeric regions and thus forming a large inverted repeat in  
483 most of the centromeres. Similar organization and specific orientation of  
484 inverted repeats also appears flanking the core centromeres of *S. pombe*  
485 [51,52] and *Candida tropicalis* [40], performing an important role in  
486 centromere function [40,53]. *M. circinelloides* is the first organism to

487 display an enrichment in retrotransposable elements in their centromeric  
488 regions in the absence of CENP-A, suggesting that these non-LTR  
489 elements determine the centromere identity. Thus, we hypothesize that  
490 identifying Grem-LINE1 clusters in other species of Mucoromycotina could  
491 help mapping the location of the putative pericentromeric regions of the  
492 corresponding species.

493         RNAi machinery is essential to control the mobility of transposable  
494 elements, but also to maintain centromere structure and stability [23].  
495 Fungi that have lost essential components of the RNAi machinery have  
496 shorter pericentric regions and fewer transposable elements than closely  
497 related RNAi-proficient species. *M. circinelloides* has a functional and  
498 complex RNAi system with Dicer-dependent and -independent pathways  
499 [24]. Our results demonstrate that the expression of the centromeric  
500 Grem-LINE1s is suppressed by the Dicer-dependent RNAi pathway, which  
501 may have resulted in accumulation of these elements in the centromeres  
502 [23].

503         Overall this study has uncovered for the first time the centromeres  
504 of an early-diverging fungus, which are characterized by a mosaic of  
505 features from both point and regional centromeres found in the fungal  
506 kingdom (Figure 6). Several studies on rapidly-evolving centromeres  
507 suggest that point centromeres may be a recently evolved state as  
508 compared to regional centromeres [9]. Therefore, we hypothesize that the  
509 last mucoralean common ancestor had regional centromeres with  
510 pericentric regions that harbored retrotransposable elements contained

511 by an active RNAi machinery. The loss of CENP-A triggered an  
512 evolutionary rearrangement from typical regional centromeres to the  
513 smaller kinetochore-binding regions that defines the mosaic centromeres  
514 of *M. circinelloides*.

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545 meetings.

## 546 **Author Contributions**

547 KS, JH, VG, and FEN conceived and supervised the study. MINM and  
548 CPA analyzed the distribution of kinetochore proteins in fungi, generated  
549 and validated the plasmids and strains, performed the ChIP-seq  
550 experiments, identified and defined the retrotransposable elements, and  
551 analyzed the mRNA and sRNA-seq data. MINM, CPA and PG analyzed the  
552 ChIP-seq data. MINM, CPA, PG and SP characterized the centromeric loci.  
553 SP, MINM and CPA did the fluorescence microscopy imaging. SP performed  
554 the ChIP-qPCR experiments. IVG, JP and SJM provided the genome  
555 assembly and gene annotation. MINM, CPA and SP designed the figures  
556 and wrote the manuscript with support from KS, JH, and VG. VG, JH, FEN  
557 and KS provided funding. All authors revised the manuscript and approved  
558 the final version.



## 559 Declaration of Interest

560 The authors declare no competing interests.

## 561 Main-text figure legends

562 **Figure 1. Distribution of the kinetochore complex across the**  
563 **Mucoromycotina subphylum.** Concise kinetochore schematic showing  
564 the most conserved protein complexes in eukaryotes is shown on the left  
565 corresponding to the kinetochore proteins analyzed in the matrix on the  
566 right. The matrix displays the presence or absence of 26 kinetochore  
567 proteins across a cladogram of 51 fungal lineages (top) to show the  
568 relationships among them. Every fungal phylum is represented and color-  
569 coded, differentiating the Mucoromycota into the Glomeromycotina,  
570 Mortierellomycotina, and Mucoromycotina subphyla to provide a special  
571 emphasis on the latter. Arrows at divergence events in the cladogram  
572 mark the hypothetical loss of proteins CENP-A (red) and CENP-C (blue) in  
573 these clades. See also Figure S1 and Table S1.

574 **Figure 2. Inner and outer kinetochore proteins colocalize in *M.***  
575 ***circinelloides* nuclei in the absence of CENP-A and CENP-C. (A)**  
576 Graphical representation of Mis12, Dsn1, and CENP-T sequence features  
577 showing individual Pfam domains PF05859, PF08202, PF15511  
578 respectively identified in *M. circinelloides*. **(B)** Schematic to represent C-  
579 terminal tagging of histone H3 and kinetochore proteins with mCherry  
580 (red circles), eGFP (green circles). **(C-F)** Confocal microscope imaging  
581 showing the cellular localization of well-conserved mCherry-tagged outer  
582 kinetochore proteins Mis12 **(C, E)** and Dsn1 **(D, F)**, eGFP-tagged histone

583 H3 **(C, D)**, and eGFP-tagged inner kinetochore CENP-T **(E, F)** in pre-  
584 germinated spores of *M. circinelloides* strains expressing fluorescent  
585 fusion proteins. **(G)** Time-lapsed confocal image displaying the cellular  
586 localization of mCherry-tagged Mis12 and eGFP-tagged histone H3 in a  
587 germinative tube sprouting from a spore. A discontinuous white perimeter  
588 outlines the germinative tube in the fluorescence images, and a white  
589 square indicates a nuclear division event. Cartoon for the zoomed image  
590 represents the localization and signal intensity of Mis12-mCherry in the  
591 nucleus. A calibrated scale (white bar) is provided for size comparison (5  
592  $\mu\text{m}$ ) and mCherry fluorescent signal is colored as magenta in the merged  
593 images in **C-G**. See also Figures S2 and S3, and Table S2.

594 **Figure 3. *M. circinelloides* displays nine centromeres.** **(A)** Putative  
595 kinetochore-binding regions showing color-coded enrichment of  
596 immunoprecipitated DNA (*IP* DNA) from Mis12 and Dsn1 mCherry-tagged  
597 strains compared to their corresponding input (*Input* DNA) and binding  
598 controls (*Beads only*). 1 kb flanking sequences from the center of the  
599 enrichment peak are shown. Major (M) and minor (m) kinetochore-binding  
600 regions are indicated. Each kinetochore-protein data was obtained from a  
601 pool of duplicated IP DNA samples. **(B)** Scaled ideogram of the nine  
602 scaffolds (white) of *M. circinelloides* genome assembly that contain a  
603 kinetochore-binding region (blue), showing the telomeric repeats (red).  
604 See also Figure S4 and Table S3.

605 **Figure 4. The centromeres of *M. circinelloides* are short, AT rich**  
606 **and harbor a conserved DNA motif.** **(A)** Size distribution of the nine

607 core centromere (CC) regions. **(B)** GC content across the nine core  
608 centromere (CC) sequences. The median (red line) and standard deviation  
609 (black lines) are shown in **A** and **B**. **(C)** 41 bp centromere-specific DNA  
610 motif conserved in all nine CC regions and absent in the remainder of the  
611 genome. A score of 2 bits in the logo indicates that the nucleotide is  
612 present at that position in all nine CC regions. **(D)** A genomic view of  
613 *CEN11* illustrating all nine core centromere regions. Kinetochore-binding  
614 region enrichment (IP coverage) as the average of both  
615 immunoprecipitation signals minus *Input* and *beads only* controls (left  
616 axis, blue), GC content across the region (right axis, black), and the  
617 centromere-specific DNA motif described in **e** (red) are shown. See also  
618 Figure S4 and Table S3.

619 **Figure 5. *M. circinelloides* core centromeres are present in large**  
620 **ORF-free pericentric regions having retrotransposable elements**  
621 **regulated by the Dicer-dependent RNAi pathway and bind H3**  
622 **nucleosomes. (A)** Genomic sequences lacking annotated genes (light  
623 blue) that flank the CC regions (dark blue) served as the reference points  
624 to calculate the length both upstream and downstream. Black circles at  
625 either end of the regions indicate an abrupt, non-telomeric end of scaffold.  
626 **(B)** Schematic of the Grem-LINE1 interspersed in the pericentromeric  
627 regions. Open ReadinE Frames (ORF) and protein domains predicted from  
628 their coding sequences are shown as colored boxes [Zf, zinc finger  
629 (PF00098 and PF16588); AP, AP endonuclease (PTHR22748); RVT, reverse  
630 transcriptase (PF00078); and zf-RVT, zinc-binding in reverse transcriptase  
631 (PF13966)]. **(C)** *CEN4* is exemplified by low GC content, lack of genes and

transcripts, which are shown in different tracks displaying the kinetochore-binding region enrichment (IP, an average of both *IP* DNA signals minus *Input* and *beads only* controls), annotation of genes (red blocks) and transposable elements (light blue blocks), *CEN*-specific DNA motif position (vertical line) and direction (arrow), GC content, and transcriptomic data of mRNA (green) and sRNAs (red) in *M. circinelloides* wild-type, *ago1*, and double *dcl1 dcl2* deletion mutant strains after 48 h of growth in rich media. **(D)** Genomic location of *CEN2* showing the regions studied for histone H3 occupancy as labelled and colored rectangles. ChIP assays were performed using polyclonal antibodies against histone H3. Primers were designed for *CC2* region, pericentric regions (1L, 2L, 1R, 2R), flanking ORFs (ORF-L, ORF-R) and a far-*CEN* control ORF ~2 Mb away from *CEN2* (Far-*CEN*). *IP* samples were analyzed by real-time PCR using these primers. The y-axis denotes the qPCR value as a percentage of the total chromatin input with standard error mean (SEM), from each region tested. The experiment was repeated three times with similar results. See also Figures S5 and S6, and Tables S2, S3 and S4.

**Figure 6. *M. circinelloides* possesses a mosaic of point and regional centromeres.** Length plot of the core centromere (kinetochore-bound region, blue circles) and the full centromere (defined as the region required for centromere function, red dots) of representative fungi belonging to the Mucoromycota (green panel), Ascomycota (blue panel) and Basidiomycota (purple panel). The size axis is divided in short ( $\leq 1$  kb, gray area) and long regions ( $> 1$  kb, brown area). The centromeres of *M. circinelloides* (CEN5), *S. cerevisiae* (CEN3 [7]), *Candida albicans* (CEN7

657 [39]), *C. tropicalis* (Scnt 3 [40]), *Magnaporthe oryzae* Guy11 (CEN2 [49]),  
658 *S. pombe* (CEN2 [52]), and *C. neoformans* (CEN10 [23]) are used for  
659 comparison and classified as point, regional and mosaic centromeres  
660 according to their structural features. In organisms where functionality of  
661 the centromere regions could not be experimentally determined, the ORF-  
662 free region spanning the kinetochore binding region is considered as the  
663 full centromere. Line diagrams represent the structural features of each  
664 centromere (drawn to scale). The ORF-free region at the centromere is  
665 defined by black vertical boundaries and flanking ORFs are shown as red  
666 arrows. Core centromeres (rectangles); kinetochore-bound regions (blue  
667 area); centromere-specific DNA motifs next to an AT-rich core (green  
668 lines); Grem-LINE1, Grasshopper, GYMAG, Maggy, Tcn retrotransposons  
669 (black arrows); inverted repeats (purple arrows); and other DNA repeats  
670 (light and dark orange arrows) are shown. Repeats/retrotransposons sizes  
671 are not drawn to scale. CC, core centromere; CDE, centromere DNA  
672 elements; LR, left repeat; RR, right repeat; otr, outer repeats containing  
673 dg/dh elements; imr, innermost repeats; cnt, central core.

## 674 **STAR Methods**

### 675 **Lead contact and materials availability**

676 Further information and requests for resources and reagents should  
677 be directed to and will be fulfilled by the Lead Contact, Victoriano Garre  
678 (vgarre@um.es).

### 679 **Experimental model and subject details**

## 680 **Fungal strains and culture conditions**

681 All the strains used and generated in this work derive from *M.*  
682 *circinelloides* f. *lusitanicus* CBS277.49. The double auxotroph MU402 [14]  
683 (Ura<sup>-</sup>, Leu<sup>-</sup>) or the MU636 (Ura<sup>+</sup>, Leu<sup>-</sup>) strain was used for transformation  
684 with cassettes containing the selectable marker for uracil *pyrG* or for  
685 leucine *leuA*, respectively. Single tagged strains in kinetochore proteins  
686 MU840, MU842, MU844, and MU846 were used as parental strains for  
687 transformation with the Hht4-eGFP cassette to obtain double-tagged  
688 strains. All the generated strains in this work are listed in the Key  
689 Resources Table.

690 After transformation, to select uracil prototrophy *M. circinelloides*  
691 spores were plated in minimal medium with Casamino Acids (MMC) [14],  
692 and for leucine prototrophy spores were plated in medium yeast nitrogen  
693 base (YNB) [14], both adjusted at pH 3.2 for transformation and colony  
694 isolation. *M. circinelloides* spores were cultured in rich medium yeast-  
695 peptone-glucose (YPG)[14] for optimal growth and sporulation at pH 4.5.  
696 The microscopy and ChIP assays were performed with pre-germinated  
697 spores growing in YPG pH 4.5 for 3 hours at 250 rpm and 26 °C.

## 698 **Method details**

### 699 **Construction of the fluorescent strains**

700 Alleles containing fluorescent fusion protein coding-sequences were  
701 designed to either integrate them in their corresponding endogenous  
702 locus, or in the *carRP* locus. The Hht4-eGFP, Hhf1-eGFP, Hhf4-eGFP and

703 CENP-T-eGFP alleles contained the ORF of each gene fused in-frame at  
704 their C-termini with the eGFP sequence, followed by the *leuA* gene as a  
705 selectable marker. The alleles were obtained by overlapping PCR using 5'-  
706 modified primers harboring restriction sites for easy cloning. Briefly, each  
707 allele was obtained by fusing the 1-kb fragment upstream the stop codon  
708 of each gene, the 0.7-kb eGFP fragment (ending in a stop codon), the 3.4-  
709 kb *leuA* fragment, and the 1-kb sequence downstream the stop codon of  
710 each tagged gene. All the linear alleles were digested with appropriate  
711 restriction enzymes, ligated into the plasmid Bluescript SK (+), and cloned  
712 into *Escherichia coli*.

713       The Mis12-mCherry and Dsn1-mCherry constructions were designed  
714 to allow the integration and ectopic overexpression of the fused proteins  
715 in the *carRP* locus. First, a universal mCherry-expression vector was  
716 constructed, containing the strong promoter *Pzrt1*, the ORF of the  
717 mCherry coding-gene including the stop codon, and the *pyrG* gene as a  
718 selectable marker (named pMAT1915). To do that, a fragment containing  
719 the 1-kb downstream and upstream sequences from the *carRP* gene  
720 flanking the *Pzrt1* promoter was amplified with an inverse PCR using the  
721 plasmid pMAT1477 [14] as a template. Then, the *mCherry* and *pyrG*  
722 fragments were added by In-Fusion cloning (Takara) using primers with  
723 appropriate overlapping 5'-tails (Table S2). Once the universal pMAT1915  
724 vector was constructed, it was used for the C-terminal tagging of *mis12* or  
725 *dsn1*, fusing their ORFs without the stop codon upstream and in-frame  
726 with the *mCherry* ORF by In-Fusion cloning. The resulting plasmids were  
727 pMAT1917 and pMAT1919, respectively for *mis12* and *dsn1*. For the N-

728 terminal tagging, the plasmid pMAT1915 was inverse-amplified using  
729 primers that exclude the stop codon of the *mCherry* fragment, and the  
730 *mis12* and *dsn1* ORFs, including their stop codons, were cloned in-frame.  
731 The resulting plasmids were pMAT1916 and pMAT1918 for mCherry-Mis12  
732 and mCherry-Dsn1 expression, respectively.

733 Stellar<sup>TM</sup> competent cells (Takara) were used for transformation by  
734 heat-shock following the supplier procedures. The correct construction of  
735 all the generated plasmids was confirmed by restriction analysis and  
736 Sanger sequencing.

737 For transformation, the alleles were released from the vector  
738 backbone by restriction digestion and used to transform *M. circinelloides*  
739 protoplasts by electroporation [14]. Spores from individual colonies were  
740 collected and plated in selective media to ensure homokaryosis. After at  
741 least five vegetative cycles, the integration in the target locus was  
742 confirmed by PCR using primers that amplified the whole locus (Table S2),  
743 producing PCR-fragments that allowed discrimination between the mutant  
744 and the wild-type locus (Figures S3). Genomic DNA purification was  
745 performed following the procedure previously published [54].

## 746 **Microscopy imaging and acquisition**

747 Freshly isolated *M. circinelloides* spores were washed twice with 1X  
748 phosphate buffered saline (PBS). The spores were resuspended in PBS and  
749 10 µl suspension was placed on a slide, covered with a coverslip and  
750 sealed with clear nail polish. The images were acquired at room  
751 temperature using laser scanning inverted confocal microscope LSM 880-



752 Airyscan (Zeiss, Plan Apochromat 63x, NA oil 1.4) equipped with highly  
753 sensitive photo-detectors. The filters used were GFP/FITC 488, mCherry  
754 561 for excitation and GFP/FITC 500/550 band pass, mCherry 565/650  
755 long pass for emission. Z- stack images were taken at every 0.5  $\mu\text{m}$  and  
756 processed using Zeiss Zen software. All the images were displayed after  
757 the maximum intensity projection of images using Zeiss Zen software.

## 758 **Live cell imaging**

759 Freshly isolated *M. circinelloides* spores were incubated in YPG  
760 growth medium at 26°C for 2 h. Then, the spores to be imaged were  
761 pelleted at 3000 xg and washed twice with 1X phosphate buffered saline  
762 (PBS). The spores were resuspended in PBS and 10 $\mu\text{l}$  suspension was  
763 placed on a slide containing thin 2% agarose patch containing 2%  
764 dextrose and covered with a coverslip. Live cell imaging was performed at  
765 30°C in a temperature-controlled chamber of an inverted confocal  
766 microscope (Zeiss, LSM-880) using a Plan Apochromat 100X NA oil 1.4  
767 objective and GaAsp photodetectors. Images were collected at 60-s  
768 intervals with 0.5  $\mu\text{m}$  Z-steps using GFP/FITC 488, mCherry 561 for  
769 excitation. GFP/FITC 500/550 band pass or GFP 495/500 and mCherry  
770 565/650 or mCherry 580-750 band pass for emission. All the images were  
771 displayed after the maximum intensity projection of images at each time  
772 using Zeiss Zen software.

## 773 **Ortholog search**

774 Seventy five fungal proteomes were retrieved from the Joint  
775 Genome Institute (JGI) Mycocosm genome portal [55] (Table S1). BLAST+

776 [56] v2.7.1 individual protein-protein BLASTp, iterative BLASTp (PSI-  
 777 BLAST) and iterative HMMER v3.2.1 (<http://www.hmmer.org>) jackhammer  
 778 searches ( $E\text{-value} \leq 1 \times 10^{-5}$ ) were conducted against these proteomes  
 779 using *M. musculus*, *U. maydis*, *S. cerevisiae*, and *S. pombe* protein  
 780 sequences (Table S1) as queries. In addition, an hmmsearch was launched  
 781 using the Pfam-A [57] HMM models for each kinetochore protein [Pfam  
 782 gathering threshold (GA) as cut-off value]. All the putative orthologs found  
 783 were used to perform a reciprocal BLASTp against *M. musculus*, *U.*  
 784 *maydis*, *S. cerevisiae*, and *S. pombe* protein sequences. Also, Pfam-A  
 785 protein domains were searched in all the possible orthologs using HMMER  
 786 hmmscan. Protein sequences that either lacked appropriate Pfam domains  
 787 or failed to produce a hit in a reciprocal blastp search were discarded,  
 788 resulting in a matrix of putative orthologs for each given species (Table  
 789 S1). The cladogram showing the relationship among species was drawn  
 790 using the tree data from JGI Mycocosm [55] and the interactive Tree of  
 791 Life[58] (iTOL v4) tool. To determine CENP-A presence or absence in each  
 792 given species, all 289 protein sequences of histone H3 putative orthologs  
 793 (Table S1) were aligned using MUSCLE[59] v3.8.1551, and the  
 794 phylogenetic relationship of their HFD was inferred by the neighbor-joining  
 795 method employing the Jones, Taylor, and Thornton (JTT) substitution  
 796 model [60], with a bootstrap procedure of 1,000 iterations (MEGA[61]  
 797 v10.0.5).

## 798 **Chromatin immunoprecipitation and sequencing**

799       The ChIP protocol for *M. circinelloides* was adapted from previous  
800 studies in other fungal models [23]. Briefly, spores from the strains MU842  
801 (expressing Mi12-mCherry) and MU846 (expressing Dsn1-mCherry) were  
802 used for independent IP experiments.  $10^7$  spores/mL of these strains were  
803 germinated in 100 ml of liquid YPG pH 4.5 medium, shaking at 250 rpm at  
804 26°C for 3 hours. Then, cells were fixed by adding formaldehyde at a final  
805 concentration of 1% for 20 minutes. The fixation was quenched by adding  
806 glycine to a final concentration of 135 mM. Cells were pelleted and ground  
807 with liquid nitrogen, and 150 mg of ground pellet was resuspended in ChIP  
808 lysis Buffer (50 mM HEPES pH 7.4, 150 mM NaCl, 1% Triton X-100, 0.1%  
809 DOC, and 1 mM EDTA), with a protease inhibitor cocktail. The sample was  
810 sonicated using a Bioruptor (Diagenode) with pulses of 30s ON/OFF for 50  
811 cycles to obtain a chromatin fragmentation between 100 and 300 bp.

812       The lysates were clarified by centrifugation at 12,000 xg for 10 min  
813 at 4°C. At this step, 100 µL were frozen at -20°C for Input DNA control.  
814 The samples were then divided into 450 µL for the antibody  
815 immunoprecipitation and 450 µL for the mock control. For IP 25 µL of Red  
816 Fluorescent Protein (RFP)-Trap Magnetic Agarose (MA) beads (Chromotek)  
817 were added and 25 µL of MA beads (Chromotek) to the binding control.  
818 Samples were incubated at 4°C overnight. After incubation, the beads  
819 were washed twice with 1 mL of the low salt wash buffer (20 mM Tris pH  
820 8.0, 150 mM NaCl, 0.1% SDS, 1 % Triton X-100, and 2 mM EDTA), twice  
821 with 1 mL of the high salt wash buffer (20 mM Tris pH 8.0, 500 mM NaCl,  
822 0.1% SDS, 1% Triton X-100 and 2 mM EDTA), once with the LiCl wash  
823 buffer (10 mM Tris pH 8.0, 1 mM EDTA, 0.25 M LiCl, 1% NP40, and 1%

824 DOC), and finally once with TE (10mM Tris pH 8.0, and 1 mM EDTA). To  
825 elute the DNA, 250 µL of TES (50 mM Tris pH 8.0, 10 mM EDTA, 1% SDS)  
826 was added to each sample and incubated for 10 min at 65°C, twice. The  
827 immunoprecipitated samples and the Input controls were de-crosslinked  
828 by incubating at 65°C overnight. The samples were treated with 199 µg of  
829 RNase A (Sigma Aldrich) and 190 µg of Proteinase K (Sigma Aldrich) for 2  
830 hours at 50°C. For DNA purification, 1 volume of  
831 phenol:chloroform:isoamyl alcohol (25:24:1) was added, the aqueous  
832 phase was recovered, and then 1 volume of chloroform:isoamyl alcohol  
833 (24:1) was added. After centrifugation, the aqueous phase was recovered,  
834 and DNA was precipitated with 20 µg of glycogen (Thermo Fisher), 1/10  
835 volume of Na-Acetate 3 M (pH 5.2), and 1 volume of ethanol 100%. Purified  
836 DNA from duplicated samples were precipitated together for sequencing.  
837 Samples incubated at -20°C overnight were centrifuged for 10 min at  
838 12,000 xg and the pellets were washed with ethanol 70%. Each sample  
839 was resuspended in 30 µL of MilliQ water. The quality of the DNA was  
840 analyzed by QuBit (Thermo Fisher) and sent to Novogene, which prepared  
841 the libraries using TruSeq ChIP Library Prep Kit and sequenced them with  
842 Illumina Hiseq 2500 High-Output v4 single-end reads of 50 bp.

843 Histone H3-ChIP assays were performed in the same way with the  
844 following changes. The protoplasts were obtained as described before [14]  
845 and then sonicated as above. Native ChIP was performed [23] using anti-  
846 H3 antibody (Abcam ab1791). The DNA pellet was dissolved in 25 µl of  
847 MilliQ water. All samples (*Input*, *IP* with or without antibodies) were used  
848 for PCR. The *Input* and *IP* DNA was subsequently used for qPCR using the

849 primers listed in Table S2 and SensiFAST™ SYBR® No ROX Kit. Histone H3  
850 enrichment was determined by the percentage input method using the  
851 formula:  $100 \times 2^{\text{adjusted Ct Input} - \text{adjusted Ct IP}}$ . The adjusted Ct value is the dilution  
852 factor ( $\log_2$  of dilution factor) subtracted from the Ct value of the *Input* or  
853 *IP* DNA samples. Three technical replicates were taken for qPCR analysis  
854 and standard error of mean was calculated.

## 855 **Sequencing data analysis**

856 Raw reads from Chromatin Immunoprecipitation and RNA  
857 sequencing (ChIP- and RNA-seq) were quality-checked with FASTQC  
858 v0.11.8 and adapters removed with Trim Galore! v0.6.2  
859 (<http://www.bioinformatics.babraham.ac.uk/projects/>). Reads were aligned  
860 to the *Mucor circinelloides* f. *Iusitanicus* MU402 reference genome  
861 Muccir1\_3 (referred to as the *M. circinelloides* genome throughout the  
862 text, available at JGI  
863 [http://genome.jgi.doe.gov/Muccir1\\_3/Muccir1\\_3.home.html](http://genome.jgi.doe.gov/Muccir1_3/Muccir1_3.home.html)), using the  
864 Burrows-Wheeler Aligner [62] (BWA v0.7.8) algorithm for ChIP- and sRNA-  
865 seq reads and STAR [63] v2.7.1a for mRNA-seq data. The number of  
866 overlapping aligned reads per 25-bp bin was used as a measure of  
867 coverage, obtaining normalized bigWig coverage files with deepTools [64]  
868 v3.2.1 bamCoverage function. For RNA-seq data, coverage was  
869 normalized to bins per million mapped reads (BPM). For ChIP-seq data,  
870 reads were normalized per genomic content (1x normalization). ChIP  
871 enrichment peaks were identified by Model-based Analysis of ChIP-seq  
872 [65] (MACS2 v2.1.1) callpeak function and sorted by fold-enrichment and

873 FDR values (Table S3). Enriched peaks (fold-enrichment  $\geq 1.6$ , FDR  $\leq$   
874  $5 \times 10^{-5}$ ) were manually curated by analyzing the *IP* coverage with the  
875 Integrative Genomics Viewer [66] (IGV v2.4.1) genome browser, ensuring  
876 that peaks were present in both Mis12 and Dsn1 *IP* samples, and absent in  
877 either the input and binding controls. The resulting centromeric sequences  
878 were analyzed with Multiple Em for Motif Elicitation [67] (MEME v5.0.5) to  
879 discover conserved DNA motifs on both strands. GC content every 25-bp  
880 bin was calculated with bedTools [68] v2.28. Normalized coverage, GC  
881 content and genetic elements annotation files were rendered with either  
882 Gviz [69] v1.29.0 for whole-chromosome visualization or deepTools  
883 pyGenomeTracks module for close-in genomic views.

#### 884 **Transposable element analysis**

885 The *M. circinelloides* genome assembly of strain MU402 was  
886 searched for repeats using RepeatScout [70] v1.0.5 and RepeatMasker  
887 (<http://www.repeatmasker.org>) v4.0.9. The repeats located in the  
888 pericentromeric regions were aligned using MAFFT [71] v7.407 and  
889 clustered with CD-HIT [72] v4.8.1, generating groups of sequences that  
890 shared  $\geq 95\%$  identity and  $\geq 70\%$  coverage and calculating their *p*-  
891 distance (nucleotide changes per site). A BLASTn search against *M.*  
892 *circinelloides* genome assembly was conducted using a representative  
893 repeated sequence from each cluster to locate similar sequences across  
894 the entire genome, obtaining hits with  $\geq 80\%$  identity and  $\geq 20\%$   
895 coverage (Table S4). The pericentromeric repeats were examined with  
896 getorf from Emboss [73] v6.6.0, obtaining all ORFs  $\geq 600$  nt. Protein

domains in these ORFs translated sequences were predicted by  
hmmsearch using the Pfam-A v32.0 database. The reverse transcriptase  
(RVT) domains found in the repeated sequences were aligned with  
MUSCLE to create a consensus sequence, and this consensus sequence  
was aligned with 281 RVT domains of well-known non-LTR  
retrotransposons from Repbase [74] 24.04. A neighbor-joining  
phylogenetic tree was inferred from this alignment using the JTT  
substitution model and a bootstrap procedure of 1000 iterations (MEGA  
v10.0.5). To assess the presence of the retrotransposable elements in  
other species of Mucoromycotina, a tBLASTn search was launched against  
the 55 available genomes at JGI (Table S5) using the translated sequence  
of both ORFs combined, recovering hits with  $\geq 50\%$  coverage and E-value  
 $\leq 10^{-5}$ .

## **Quantification and statistical analysis**

Statistical details of experiments are detailed in the Results, Figures  
and Figure legends, including the number of biological and technical  
replicates, the definition of center, dispersion and precision measures  
(mean, SEM and SD). Statistical analyses were performed with GraphPad  
Prism 6.

## **Data and code availability**

ChIP-seq raw data and processed files are deposited at the Gene  
Expression Omnibus (GEO) repository under GSE132687 accession  
number and are publicly available. PacBio raw data and assembly of the

920 genome of *M. circinelloides* f. *lusitanicus* MU402 are publicly available at  
921 the JGI website  
922 ([http://genome.jgi.doe.gov/Muccir1\\_3/Muccir1\\_3.download.html](http://genome.jgi.doe.gov/Muccir1_3/Muccir1_3.download.html)) and  
923 subjected to the JGI Data Usage Policy. Both mRNA and sRNA-seq data  
924 were already available at the Sequences Read Archive (SRA) and can be  
925 accessed with the following run accession numbers: SRR1611144 (R7B  
926 wild-type strain mRNA) [75], SRR1611151 (*ago1* deletion mutant strain  
927 mRNA) [75], SRR1611171 (double *dcl1 dcl2* deletion mutant strain mRNA)  
928 [75], SRR039123 (R7B wild-type strain sRNA) [76], SRR836082 (*ago1*  
929 deletion mutant strain sRNA) [77], SRR039128 (double *dcl1 dcl2* deletion  
930 mutant strain sRNA) [76]. The protein sequences of the kinetochore  
931 orthologs identified in this study are accessible at Mendeley data (DOI:  
932 10.17632/wpyyb58h8v.1), as well as all the main and supplementary  
933 material.

#### 934 **Legends for supplemental Excel tables and videos**

935 **Video S1. *M. circinelloides* displays asynchronous nuclear**  
936 **division. Related to Figure 2.** A pre-germinated spore of *M.*  
937 *circinelloides* showing the distribution of histone H3-eGFP fusion protein  
938 encompassing the nuclei. Red arrows mark single events of nuclear  
939 division.

940 **Table S1. Kinetochore proteins distribution across fungal**  
941 **lineages. Related to Figure 1.** Orthologs identified in each species  
942 analyzed, showing their correspondent ID in the JGI platform, Pfam



domains (name, ID, coordinates within the translated sequence, e-value and score).

**Table S3. Centromere location and Mis12 and Dsn1 IP/Input peaks. Related to Figures 3, 4, 5 and S4. (A)** Genomic coordinates of the core centromere (CC) defined by the both Mis12 and Dsn1 kinetochore proteins binding regions; and of the entire centromere, including the upstream and downstream pericentric regions. **(B, C)** MACS2 called peaks for Mis12 IP/Input **(B)** and Dsn1 IP/Input **(C)**. Rows colored red are above established threshold ( $\text{FDR} \leq 5 \times 10^{-7}$ , fold enrichment  $\geq 1.6$ ).

**Table S4. Centromeric Grem-LINE1s domain architecture. Related to Figures 5, S5 and S6.** Protein domains [name and ID (Pfam or Panther), start and end in the translated product, as well as predicted e-value, score, and bias] for the ORFs [numbered (#) and showing start and end in the DNA sequence] found in each Grem-LINE1 (showing genomic location and length).

**Table S5. Grem-LINE1s distribution across Mucoromycotina. Related to Figures 1 and S6.** Genomic location of Grem-LINE1s across species of Mucoromycotina showing the percentage of identity compared to a representative of *M. circinelloides* Grem-LINE1s.

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